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Particule de percarbonate de sodium stable et procédé de préparation

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Description

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Background of the Invention

This invention relates to a sodium percarbonate particle having excellent stability and also to a process for preparing the same.

Sodium percarbonate is well-known as a bleaching or oxidizing agent. It is commonly manufactured by the reaction of sodium carbonate with hydrogen peroxide. As its general formula $2Na_2CO_33H_2O_2$ implies, it comprises an addition compound of sodium carbonate with hydrogen peroxide. Although sodium percarbonate is somewhat inferior in bleaching power at ordinary temperature to chlorinated bleaches, it can be used in bleaching textile pieces with colored patterns, does not damage the fabric, and causes no yellowing. These and other advantages have allowed it to enjoy widespread use, alone or in admixture with detergents, as oxygenated bleaching agents for household or business use.

The reason why an interest is attached to the use of sodium percarbonate in applications such as general detergents and domestic bleaches is that its decomposition products are not pollutants and can be used in whatever way desired without any problem.

However, sodium percarbonate has a serious drawback in that it is rather poor in storage stability as compared with sodium perborate and fairly rapidly loses its available oxygen during storage. Since sodium percarbonate has a very high affinity towards water, it becomes wetted on its surface even with a small amount of moisture and begins decomposing. Iron, copper, manganese, cobalt, and other such ions, if present in the system, accelerate its decomposition, making it less stable than sodium perborate. When sodium percarbonate, alone, is stored in a closed vessel, it proves to be as stable during storage as sodium perborate. When blended with a detergent or left in an open vessel, however, it exhibits inadequate storage stability because of its high hygroscopicity, despite its good solubility.

It is therefore desired to provide a sodium percarbonate which does not decompose rapidly upon standing in an open vessel and, when blended with whatever detergent or bleach composition, does not present any storage stability problems.

Many different researches have been conducted with regard to the stabilization of sodium percarbonate and various approaches have been proposed. Particularly, a large number of proposals have been made with respect to coating the surface of a sodium percarbonate particle with an inorganic matter.

For instance, Japanese Patent Publication No. 31839/1972 discloses sodium percarbonate surface-coated with sodium perpyrophosphate; U.S. Patent No. 3977988, sodium percarbonate surface-coated with a silicate-silicofluoride mixture; Japanese Patent Publication No. 5875/1978, sodium percarbonate surface-coated with a silicate-glycine mixture; U.S. Patent No. 4105827, sodium percarbonate surface-coated with a crystallized mixed salt consisting of sodium carbonate and either sodium bicarbonate or sodium sulfate; U.S. Patent No. 4194025, sodium percarbonate surface-coated with dehydrated sodium perbonate and sodium silicate; U.S. Patent No. 4325933, sodium percarbonate surface-coated with dehydrated sodium perbonate; Japanese Patent Laid-Open No. 193999/1984, sodium percarbonate surface-coated with a borate-silicate mixture; Japanese Patent Laid-Open No. 194000/1984, sodium percarbonate surface-coated with a borate-magnesium compound mixture; Japanese Patent Laid-Open No. 118606/1985, sodium percarbonate surface-coated with a mixture of boric acid or borate and a water repellent agent; and U.S. Patent No. 4526698, sodium percarbonate surface-coated with a satisfactory extent.

FR-A-2528447 discloses a one-step process in which moist sodium percarbonate particles and hydrated sodium metaborate particles are mixed together and dried.

The present inventors have made investigations in particular on the stabilization of sodium percarbonate particles. It was presumed in the course of research that, while a sodium percarbonate particle has numerous pores within the particle, the inorganic salt-coated sodium percarbonate particle of the prior art is one coated only on its surface, the coating being so imperfect as to leave minute interstices on the surface so that the particle absorbs moisture through these interstices and disintegrates from the inside thereof. However, absolutely perfect coating of the particle surface with an inorganic salt is next to impossible and, if ever possible, would require a very large amount of the inorganic salt. After further intensive research under the circumstances, the present inventors have found that a sodium percarbonate particle prepared by penetrating and coating a borate to its pores attain exceptionally good stability and have accomplished the present invention on that basis.

Accordingly the present invention provides for a stable sodium percarbonate particle which comprises a sodium percarbonate particle and a borate wherein at least the inside of said particle is penetrated and coated with said borate.

The present invention also provides for a stable sodium percarbonate particle which comprises a sodium percarbonate particle and a borate wherein the inside of said particle is penetrated and coated with said borate and the surface of said particle is coated with said borate, the amount of borate in the stable sodium percarbonate particle being from 0.5 to 10 % by weight of the sodium percarbonate.

Furtheremore the present invention provides for a process for preparing a stable sodium percarbonate particle which comprises (a) subjecting a sodium percarbonate particle and an aqueous solution of a borate to solid-liquid mixing at a temperature of 35°C or below, (b) drying the resulting mixture and (c), repeating steps (a) and (b) at least once more. Non-limiting Examples of the invention will now be describes in certain instances with reference to the accompanying drawings, in which:

Fig. 1 is a scanning electron micrograph of the structure of a sodium percarbonate particle obtained in Example 1; and Fig. 2 is a scanning electron micrograph of the structure of a sodium percarbonate particle obtained in Comparative Example 1.

A stable sodium percarbonate particle of the present invention is at least the inside of the particle is penetrated and coated with a borate. In other words, a stable sodium percarbonate particle of the present invention is a sodium percarbonate composite comprising sodium percarbonate particle and a borate with which the walls of pores existing in a sodium percarbonate particle are covered and coated.

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The borate according to the invention is preferably a sodium salt of boric acid, especially preferably sodium metaborate. A sequestering agent, such as ethylenediaminetetraacetate, nitrilotriacetate and a phosphate, and an inorganic salt such as a silicate and a sulfate may be contained in the sodium percarbonate composite together with the borate.

The amount of the borate is from 0.5 to 10% by weight, preferably from 2 to 8% by weight, based on the weight of sodium percarbonate.

The sodium percarbonate composite desirably has an average particle diameter of from 100 to 2000 μ m, more desirably from 250 to 1500 μ m.

Namely the sodium percarbonate particle of the present invention is obtained by treating sodium percarbonate particle with a solution of a borate so that the borate may penetrate into the inside of the particle and drying the resultant so that the borate is left there covering the walls' surface of the pores in the particle.

A sodium percarbonate obtained by reacting sodium carbonate with hydrogen peroxide in the usual manner and dehydrating the reaction product by the usual method normally contains from 7 to 18% by weight of water, and is known as wet sodium percarbonate. A sodium percarbonate prepared by drying the wet sodium percarbonate to a reduced water content is termed dry sodium percarbonate. As the raw material for the manufacture of a penetrated and coated, a stable sodium percarbonate particle in accordance with the present invention, it is advantageous to use a sodium percarbonate particle dried to a water content of less than 8% by weight. The drying presumably deforms the minute voids or pores to have the shape and size easily penetratable by the aqueous solution of a borate. Sodium percarbonate containing less than 8% by weight of water can be commercially obtained and the example of them includes sodium percarbonate manufactured by Nihon Peroxide Co. Ltd. and sodium percarbonate manufactured by Mitsubichi Gas Chemical Co. Ltd..

The sodium percarbonate as the raw material desirably ranges in average particle diameter from 50 to 2000 µm.

The borate used as a penetrating and coating agent according to the process for preparing a stable sodium percarbonate particle of the invention is preferably a sodium salt of boric acid such as sodium tetraborate decahydrate (or borax) (Na₂O·2B₂O₃·10H₂O or Na₂B₄O₇·10H₂O), sodium tetraborate pentahydrate (Na₂O·2B₂O₃·5H₂O), sodium tetraborate tetrahydrate (Na₂O·2B₂O₃·4H₂O), sodium tetraborate anhydrate (Na₂O·2B₂O₃), sodium octaborate tetrahydrate (Na₂O·4B₂O₃·4H₂O), sodium pentaborate pentahydrate (Na₂O·5B₂O₃·10H₂O), sodium metaborate tetrahydrate (Na₂O·4B₂O) and sodium metaborate dihydrate (Na₂O·2H₂O), especially preferably sodium metaborate dihydrate and sodium metaborate tetrahydrate. This agent may contain a sequestering agent, such as ethylenediaminetetraacetate, nitrilotriacetate and a phosphate. It may further contain an inorganic salt such as a silicate and a sulfate. The amount of these additives is preferably 50% and less by weight on the basis of the borate.

The amount of the borate required for penetration and coating of a sodium percarbonate particle is from 0.5 to 10% by weight, preferably from 2 to 8% by weight, based on the weight of sodium percarbonate.

The concentration of the borate in aqueous solution as the penetrating and coating agent ranges desirably from 1 to 55% by weight as anhydrous salt, more desirably from 5 to 50% by weight as anhydrous salt and especially desirably from 20 to 50% by weight as anhydrous salt.

In order to obtain a sodium percarbonate particle penetrated and coated inside and coated on the surface thereof with a borate, it is necessary to choose a proper solid-liquid mixing temperature depending on the properties of the sodium percarbonate particle used as a raw material and the type and concentration of the borate used.

It is necessary to repeat the solid-liquid mixing and drying procedures at least twice, preferably two or three times. While the solid-liquid mixing and drying are conducted at the same time, the objective stable sodium percarbonate particle of the invention may not be obtained.

The solid-liquid mixing is performed at a temperature of 35°C or below. If the particle temperature is higher than 50°C, the penetrating and coating agent fails to penetrate satisfactorily upon contact with the particle, because the dissolution rate of the particle surface exceeds the penetration rate of the penetrating and coating agent, resulting in a reduced stability of the sodium percarbonate particle in the invention. At the same time, the penetrating and coating

agent tends to act as a binder to thereby increase the average particle diameter of the coated particle above the size desired for the purposes of the invention, thus lowering the solubility of the product.

The sodium percarbonate particle penetrated and coated with a borate in conformity with the invention is by far the more stable than the conventional sodium percarbonate particle coated with a borate merely on the surface. Ordinarily, a sodium percarbonate particle with a relatively large particle diameter has been used for added stability, with consequent reduction in the solubility. According to the invention, by contrast, an increased stability is attained with a sodium percarbonate particle of a relatively small diameter. This offers an advantage from the solubility standpoint.

Example 1

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400 g of sodium percarbonate (moisture content: 0.5% by weight; average particle diameter: 495 µm) (at a solid material temperature of 20°C) was fed into an agitated mixer ("High-speed mixer FS-GS-15"; mfd. by Fukae Kogyo Co. Ltd.), While the feed was stirred at 300 rpm, 34.9 g of an aqueous solution (28.6% by weight as anhydrous salt) of sodium metaborate tetrahydrate (at a liquid temperature of 70°C) was added thereto dropwise in 3 minutes and then mixed for one minute (with circulation of water at 20°C through a jacket). Next, the mixture was transferred to a fluidizedbed ("STREA-1" type; mfd. by Powrex Co. Ltd.), where it was dried with hot air at 80°C for 10 minutes and then cooled with air at 20°C for 5 minutes. It was returned to the agitated mixer, and the above procedure was repeated. The amount of the aqueous solution containing 28.6% by weight of sodium metaborate tetrahydrate used was 21.0 g.

Example 2

The same procedure of the Example 1 was proceeded except for using an aqueous solution (19% by weight as anhydrous salt) of borax (Na₂B₄O₇·10H₂O) (at a liquid temperature of 70°C) as the penetrating and coating agent. The amount of the consumed agent was 40.0 g in the first cycle and 24.0 g in the second.

Example 3 to 5

The same procedure of the Example 1 was proceeded except that, in addition to the sodium metaborate, another inorganic salt and/or stabilizer was added to the penetrating and coating agent. The penetrating and coating agents employed, and the concentration and the amount of the aqueous solution are listed in Table 1.

The total amount of the sodium metaborate, another inorganic salt and stabilizer used was 4% by weight as anhydrous salt based on sodium percarbonate as raw material(at a liquid temperature of 70°C). The whole amount of the above-described compounds added in the first run was 2.5% by weight and that in the second run was 1.5% by weight.

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Table 1

Example No.	Penetrating and coating agent (by weight as anhydrous salt)	Amount of aqueous solution added (g)		Concentration of the aqueous solution containing penetrating and coating agent (by weight as anhydrous salt)
		first	second	
3	sodium metaborate 3% + mag- nesium sulfate 1%	33.3	20.0	sodium metaborate 22.5% + magnesium sulfate 7.5%
4	sodium metaborate 3% + sodium silicate #2 1%	33.3	20.0	sodium metaborate 22.5% + sodium silicate #2 7.5%
5	sodium metaborate 3.4% + EDTA-2Na 0.6%	36.8	22.1	sodium metaborate 23.1% + EDTA·2Na 4.1%

Comparative Example 1

The same procedure as described in the Example 1 was carried out except that the temperature of sodium percarbonate was 60°C during mixing (while hot water at 70°C was circulated through a jacket).

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Comparative Example 2

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The same procedure as described in the Example 1 was carried out except that sodium carbonate (a aqueous solution of 30% by weight as anhydrous salt) was used in place of sodium metaborate. The amount of the coating sodium carbonate was 4% by weight as anhydrous salt based on sodium percarbonate as raw material.

In the Examples 1 to 5, each sodium percarbonate particle obtained had been penetrated and coated both inside and on the outer surface of the particle. Each sodium percarbonate particle of Comparative Examples 1 and 2 was coated only on the surface of the particle.

Fig. 1 is a scanning electron micrograph (magnification 110X) of the structure of a sodium percarbonate particle obtained in Example 1. The particle is penetrated inside and coated on the surface with a metaborate. Fig. 2 is a scanning electron micrograph (magnification 110X) of the structure of a sodium percarbonate particle obtained in Comparative Example 1. The borate coating is confined to only the particle surface.

The sodium percarbonate particles prepared in the Examples and Comparative Examples as above were tested for their storage stability by the following method. The results are given in Table 2.

(Testing method)

1 g of each sodium percarbonate particles prepared in the Examples and Comparative Examples and 9 g of a phosphorus-free powdered detergent of the composition specified below were placed in a plastic container having a capacity of 50 cc. The container was closed with a lid (which had been pinholed), and the contents were allowed to stand at 40°C and 80% RH (Relative Humidity) for 14 days. The available residual oxygen rate was calculated from the following equation:

available residual oxygen rate (%) = $\frac{\text{available oxygen after storage}}{\text{available oxygen before storage}} \times 100$

For the determination of available oxygen, the method of titration with 0.1N potassium permanganate was used.

Composition of the detergent

30	polyoxyethylene (E0 = 8.5) alkyl (C_{12}) ether	17% by weight	
35	<pre>sodium linear alkyl- (C = 13.9)benzenesulfonate</pre>	3	
	beef tallow soap	2	
*	synthetic zeolite (type 4A)	39	
40	sodium silicate #1	11	
	sodium carbonate	8	
45	polyethylene glycol (MW: 14000)	2	
50	maleic acid-acrylic acid copolymer (monomer ratio: 30/70 MW: 70,000	2	
	protease	1	
	sodium sulfate	10	
55	water	5	

The solubilities of the sodium percarbonate particles obtained in the above Examples and Comparative Examples were evaluated in the following manner.

(Method of evaluating solubility)

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One liter of tap water was poured into a beaker (of 1-liter capacity), into which was thrown 1 g of each sodium percarbonate particles obtained in the above Examples and Comparative Examples, and the feed was stirred at a speed of 200 rpm. The period of time from the initiation of stirring till the point where the test solution no longer underwent a change in electric conductivity was measured and defined as the dissolution time.

Table 2 gives the results.

Table 2

Sodium percar- bonate particle	Penetrating and coat- ing agent	Avail. resid. O ₂ rate (%)	Dissolution rate (sec)	Average particle diameter (µm)
Example 1	sodium metaborate	99.8	97	- 585
Example 2	borax	98.0	98	597
Example 3	sodium metaborate + magnesium sulfate	97.8	101	603
Example 4	sodium metaborate + sodium silicate #2	99.0	105	613
Example 5	sodium metaborate + EDTA:2Na	98.5	99	600
Comp. Ex. 1	sodium metaborate	83.0	157	812
Comp. Ex. 2	sodium carbonate	46.0	132	798
Untreated sodium percarbonate		31.1	60	495

Claims

- A stable sodium percarbonate particle comprising a porous sodium percarbonate particle in which both the internal surfaces of the pores and the external surface of the particle are coated with a borate, the amount of borate in the stable sodium percarbonate particle being from 0.5 to 10% by weight of the sodium percarbonate.
- 2. The stable sodium percarbonate particle according to claim 1, wherein the borate is a sodium borate.
 - 3. The stable sodium percarbonate particle according to claims 1 or 2, wherein the borate is sodium metaborate.
 - 4. The stable sodium percarbonate particle according to claims 1 or 2, which has an average particle diameter of from 100μm to 2000μm.
 - The stable sodium percarbonate particle according to claim 4, which has an average particle diameter of from 250 μm to 1500 μm.
- 6. A process for preparing a stable sodium percarbonate particle which comprises (a) subjecting a sodium percarbonate particle and an aqueous solution of a borate to solid-liquid mixing at a temperature of 35°C or below, (b) drying the resulting mixture and (c), repeating steps (a) and (b) at least once more.
 - 7. The process according to claim 6, wherein steps (a) and (b) are repeated at least twice more.

8. The process according to claim 6, wherein the sodium percarbonate particle is dried to a water content of less than 8% by weight.

- 9. The process according to claim 6, wherein the aqueous solution of a borate contains a sequestering agent and/or an inorganic salt besides borate.
- 10. The process according to claim 6, wherein the concentration of said borate in the aqueous solution is in the range of from 1 to 55% by weight a anhydrous salt.
 - 11. The process according to claim 6, wherein said borate is used in amount of from 0.5 to 10% by weight as anhydrous salt based on sodium percarbonate as raw material.

10 Patentansprüche

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- Stabile Natriumpercarbonatteilchen umfassend poröse Natriumpercarbonatteilchen bei denen sowohl die inneren Oberflächen der Poren und die äußeren Oberflächen der Teilchen mit einem Borat beschichtet sind, wobei die Menge des Borats in den stabilen Natriumpercarbonatteilchen 0,5 bis 10 Gew.-%, bezogen auf das Natriumpercarbonat, beträgt.
- 2. Stabile Natriumpercarbonatteilchen nach Anspruch 1, worin das Borat ein Natriumborat ist.
- 3. Stabile Natriumpercarbonatteilchen nach Anspruch 1 oder 2, worin das Borat Natriummetaborat ist.
- 4. Stabile Natriumpercarbonatteilchen nach Anspruch 1 oder 2, welche einen durchschnittlichen Teilchendurchmesser von 100 bis 2000 μm besitzen.
- Stabile Natriumpercarbonatteilchen nach Anspruch 4, welche einen durchschnittlichen Teilchendurchmesser von
 250 bis 1500 μm besitzen.
 - 6. Verfahren zur Herstellung stabiler Natriumpercarbonatteilchen umfassend (a) ein Feststoff-Flüssigkeit-Mischen von Natriumpercarbonatteilchen und einer wäßrigen Boratlösung bei einer Temperatur von 35°C oder weniger, (b) Trocknen der resultierenden Mischung und (c) wenigstens einmaliges Wiederholen der Schritte (a) und (b).
 - 7. Verfahren nach Anspruch 6, worin die Schritte (a) und (b) wenigstens zweimal wiederholt werden.
 - 8. Verfahren nach Anspruch 6, worin die Natriumpercarbonatteilchen bis zu einem Wassergehalt von weniger als 8 Gew.-% getrocknet werden.
 - Verfahren nach Anspruch 6, worin die w\u00e4\u00derige Borat\u00f6\u00f3ung ein Sequestriermittel und/oder ein anorganisches Salz neben dem Borat enth\u00e4lt.
- 10. Verfahren nach Anspruch 6, worin die Konzentration des Borats in der w\u00e4\u00dfrigen L\u00f6sung im Bereich von 1 bis 55
 Gew.-%, als wasserfreies Salz, liegt.
 - 11. Verfahren nach Anspruch 6, worin das Borat in einer Menge von 0,5 bis 10 Gew.-%, als wasserfreies Salz, bezogen auf Natriumpercarbonat als Rohmaterial, verwendet wird.

5 Revendications

- Particule de percarbonate de sodium stable, comprenant une particule poreuse de percarbonate de sodium dont les surfaces internes des pores et la surface externe de la particule sont recouvertes d'un borate, la proportion de borate dans la particule de percarbonate de sodium stable étant de 0,5 à 10 % du poids du percarbonate de sodium.
- 2. Particule de percarbonate de sodium stable selon la revendication 1, dans laquelle le borate est un borate de sodium.
- 3. Particule de percarbonate de sodium stable selon la revendication 1 ou 2, dans laquelle le borate est le métaborate de sodium.
- 4. Particule de percarbonate de sodium stable selon les revendications 1 ou 2, qui a un diamètre particulaire moyen de 1 00 μm à 2 000 μm.

- 5. Particule de percarbonate de sodium stable selon la revendication 4, qui a un diamètre particulaire moyen de 250 μ m à 1 500 μ m.
- 6. Procédé pour préparer une particule de percarbonate de sodium stable, qui consiste à (a) soumettre une particule de percarbonate de sodium et une solution aqueuse d'un borate à un mélange solide-liquide à une température de 35°C ou moins, (b) sécher le mélange obtenu et (c) répéter les étapes (a) et (b) au moins encore une fois.
 - 7. Procédé selon la revendication 6, dans lequel les étapes (a) et (b) sont répétées au moins encore deux fois.
- 8. Procédé selon la revendication 6, dans lequel la particule de percarbonate de sodium est séchée jusqu'à une teneur en eau inférieure à 8 % en poids.

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- 9. Procédé selon la revendication 6, dans lequel la solution aqueuse d'un borate contient un agent séquestrant et/ou un sel minéral, en plus du borate.
- 10. Procédé selon la revendication 6, dans lequel la concentration dudit borate dans la solution aqueuse est dans la gamme de 1 à 55 % en poids en sel anhydre.
- 11. Procédé selon la revendication 6, dans lequel ledit borate est utilisé en une proportion de 0,5 à 10 % en poids en sel anhydre, relativement au percarbonate de sodium constituant la matière première.

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Fig. 1



1772 3.75% 21434 (25% (125%))

Fig. 2



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